Document made available under the Patent Cooperation Treaty (PCT)

International application number: PCT/EP05/003211

International filing date: 21 March 2005 (21.03.2005)

Document type: Certified copy of priority document

Document details: Country/Office: US

Number: 60/554,808

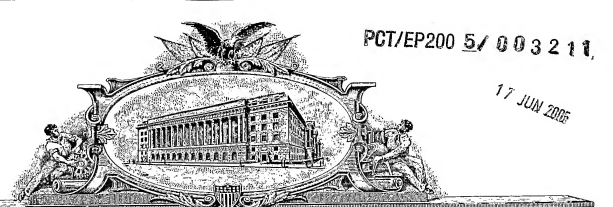
Filing date: 19 March 2004 (19.03.2004)

Date of receipt at the International Bureau: 02 August 2005 (02.08.2005)

Remark: Priority document submitted or transmitted to the International Bureau in

compliance with Rule 17.1(a) or (b)





PA 1323162

THICK OUTHRD CHARRED DEVICES OF

TO AUG TO WHOM THUSSE: PRUSDENUS SHAVE COMES

UNITED STATES DEPARTMENT OF COMMERCE

United States Patent and Trademark Office

May 27, 2005

THIS IS TO CERTIFY THAT ANNEXED HERETO IS A TRUE COPY FROM THE RECORDS OF THE UNITED STATES PATENT AND TRADEMARK OFFICE OF THOSE PAPERS OF THE BELOW IDENTIFIED PATENT APPLICATION THAT MET THE REQUIREMENTS TO BE GRANTED A FILING DATE UNDER 35 USC 111.

APPLICATION NUMBER: 60/554,808

FILING DATE: March 19, 2004

By Authority of the

COMMISSIONER OF PATENTS AND TRADEMARKS

N. WOODSON

Certifying Officer

PTO/SB/16 (08-03)
Approved for use through 07/31/2006. OMB 0651-0032
Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE
Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number.

PROVISIONAL APPLICATION FOR PATENT COVER SHEET

This is a request for filing a PROVISIONAL APPLICATION FOR PATENT under 37 CFR 1.53 (c).

INVENTOR(S)									
Given Name (first and m	Family Name o		Residence (City and either State or Foreign Country)						
Jőrn	Lewi	n	Berlin, Germany						
Additional inventors are being named on the separately num				ered sheets attached hereto					
	TITLE C	F THE INVENTION	l (500 chara	cters max)	•	2.4			
A METHOD TO ASS	ESS MEASURE	MENT METHOD	S QUANTII	FYING BASE	COMPOSITI	ONS IN DNA			
	CORRES	PONDENCE AD	DRESS			66			
Direct all correspondence to		00704	 1			"			
X Customer Number		22504							
OR	Туре	Customer Number	here						
Firm or Individual Name									
Address					<u>.</u>				
Address									
City		S	tate		ZIP				
Country		Tele	phone		Fax				
ENCLOSED APPLICATION PARTS (check all that apply) Specification Number of Pages 5 CD(s), Number Drawing(s) Number of Sheets 5 Other (specify) Fee tymsmittal Application Data Sheet. See 37 CFR 1.76 Sheet in duplicate METHOD OF PAYMENT OF FILING FEES FOR THIS PROVISIONAL APPLICATION FOR PATENT Applicant claims small entity status. See 37 CFR 1.27. A check or money order for \$ is enclosed to cover the filing fees. The Commissioner is hereby authorized to charge filing fees to Deposit Account Number: 04-0258 The Commissioner is hereby authorized to charge any deficiency or credit any overpayment to Deposit Account Number: 04-0258 Payment by credit card. Form PTO-2038 is attached. The invention was made by an agency of the United States Government or under a contract with an agency of the United States Government or under a contract with an agency of the United States Government or under a contract with an agency of the United States Government contract number are: Yes, the name of the U.S. Government agency and the Government contract number are:									
Respectfully submitted,	0	0							
SIGNATURE	Bull	Huses	DATE		March 19,	2004			
TYPED or PRINTED NAME	Bruce A. Kase	r	REGISTRA	ATION NO. iate)					
TELEPHONE	206-628-7653		DOCKET	NUMBER:	47675-				

USE ONLY FOR FILING A PROVISIONAL APPLICATION FOR PATENT

This collection of information is required by 37 CFR 1.51. The information is used by the public to file (and by the PTO to process) a provisional application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 8 hours to complete, including gathering, preparing, and submitting the complete provisional application to the PTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Mail Stop Provisional Patent Application, Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

R:\FORMS\IP Pending\Patents\PTO FORMS\SB16 Provisional App for Patent Cover Sheet.doc

Copy provided by USPTO from the IFW Image Database on 05/18/2005

EXPRESS MAIL NO. EL852794988US

PTO/SB/17 (10-03)

Approved for use through 07/31/2006. OMB 0651-0032

U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE
to a collection of information unless it displays a valid OMB control information.

Under the Paperwork Reduction Act of 1995, no persons are required to	respond to a collection of information unless it display	s a valid OMB control number.				
	Complete if Knowr	Complete if Known				
FEE TRANSMITTAL	Application Number					
	Filing Date March 19, 2004					
for FY 2004	First Named Inventor Lewin					
Effective 10/01/2003. Patent fees are subject to annual revision.	Examiner Name					
Applicant claims small entity status. See 37 CFR 1.27	Art Unit					
TOTAL AMOUNT OF PAYMENT (\$) 80	Attorney Docket No. 47675-					
	TES ON OUR ATION (or					
METHOD OF PAYMENT (check all that apply)	FEE CALCULATION (continued)					
Check Credit card Money Order None	3. ADDITIONAL FEES Large Entity Small					

METHOD OF PAYMENT (check all that apply)			FEE CALCULATION (continued)						
Check Credit card Money Order None			3. ADDITIONAL FEES Large Entity Small						
Deposit Account:			Fee Code	Fee (\$)	Fee Code	Fee (\$)	Fee Description	Fee Paid	
Deposit Account 04-0258			1051	130	2051		Surcharge - late filing fee or oath		
Number			1052	50	2052	25	Surcharge - late provisional filing fee or cover sheet.		
Deposit Account Davis Wright Tremaine LLP			1053	130	1053	130	Non-English specification		
Name			1812	2.520	1812	2,520	For filing a request for ex parte		
transmit .	r is authorized	to: (check all that appl	ly)	1804	920*	1804	920*	reexamination Requesting publication of SIR prior to	
Charge fee(s) indicated below Credit any overpayments			1004				Examiner action		
Charge any additional fee(s) during the pendency of this application			1805	1,840*	1805	1,840*	Requesting publication of SIR after Examiner action		
Charge fee(s) indicated below, except for the filing fee			1251	110	2251	55	Extension for reply within first month		
			1252	420	2252	210	Extension for reply within second month	1	
Charge any deficiencies			1253	950	2253	475	Extension for reply within third month		
to the above-Identified deposit account. FEE CALCULATION			1254	1,480	2254	740	Extension for reply within fourth month		
1. BASIC FILING FEE				1255	2,010	2255	1005	Extension for reply within fifth month	
	mall Entity			1401	330	2401	165	Notice of Appeal	
Fee	Fee			1402	330	2402	165	Filing a brief in support of an appeal	
	Code Fee(\$)	Fee Description	Fee Paid	1403	290	2403	145	Request for oral hearing	
1002 340	2002 1	85 Utility filing fee 70 Design filing fee		1451	1,510	1451	1,510	Petition to institute a public use proceeding	
1003 530		65 Plant filing fee		1452	110	2452	55	Petition to revive - unavoidable	
1004 770		85 Reissue filing fee	<u> </u>	1453	1,330	2453	665	Petition to revive – unintentional	
1005 160	2005	80 Provisional filing fee	80	1501	1,330	2501	665	Utility Issue fee (or relssue)	
· ·		SUBTOTAL (1)	(\$)80	1502	480	2502	240	9	
		30BIOIAL(I)	(4)ov	1503	640	2503	320		
2. EXTRA CLAIM	FEES			1460	130	1460	130	Petitions to the Commissioner	
		Fe Extra from	m Fee	1807	50	1807	50	Petitions related to provisional applications	
Total	- 20** =	Claims belo	ow Paid	1806	180	1806	180	Sunt	
Claims				8021	40	8021	40	Recording each patent assignment per property (times number of properties)	
Claims	-3**=	×		1809	770	2809	385		
Dependent	Complete Control		=	1810	770	2810	385		
Large Entity Fee Fee	Small Entity Fee Fe	ee (\$) Fee Description		1801	770	2801	385		
Code (\$)	code	•		1802	900	1802	900	Request for expedited examination of a	
1202 18 1201 86	2202 2201	9 Claims in excess 43 Independent clai	s of 20 ms in excess of 3	Others (- /			design application	
1203 290	2203	145 Multiple depende	ent claim, if not paid	Other fee	e (specir	у)	-		L
1204 86	2204	43 ** Reissue inder original paten	oendent claims over t	*Reduce	ed by Ba	sic Filin	g Fee P	aid SUBTOTAL (3) (\$)0	
1205 18	2205	9 ** Reissue claim over original p	ns in excess of 20 and patent						
		UBTOTAL (2)	\$)0						
**ar number proving		r; For Reissues, see above							
or nonner previou	any paru, ii greater	,, , ,, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		4					

SUBMITTED BY				(Cor	nplete (if applicable))
Name (Print Type)	Bruce A Kaser	Registration No. (Attorney/Agent)	31,531	Telepho	one 206-628-7653
Signature	Bung Cotases			Date	March 19, 2004

WARNING: Information on this form may become public. Credit card information should not be included on this form. Provide credit card information and authorization on PTO-2038.

This collection of information is required by 37 CFR 1.17 and 1.27. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14.

A method to assess measurement methods quantifying base compositions in DNA

Jörn Lewin Epigenomics AG Berlin

19th March 2004

Method

I hereby explain the method with help of an example experiment used for calibration. In the example experiment the test system was used to assess cytosine/thymine base ratio measurement methods as used in most methylation detection protocols using bisulphite treatment of the DNA.

- First identical regions with local differences are subcloned into plasmids. In the experiment this was an inhomogeneous PCR product from incomplete bisulphite treated DNA that resulted in a mixture of molecules with different C/T proportions at all positions that were cytosine prior to conversion with bisulphite.
- A set of the subclones is sequenced to obtain information about the base composition differences. Other methods to determine these differences can be used but sequencing of the subclones is the most appropriate method. In the experiment we sequenced 96 subclones from one inhomogeneous amplificate.
- A set of subclones is chosen, that compared to each other are different at as many positions as possible relevant for the measurement method to be assessed. For this method a number of two chosen subclones is the minimum but three or more lead to a higher resolution. In the experiment we chose three clones which differed at positions that in the genomic sequence were cytosine and resulted in either cytosine or thymine dependent on the bisulphite conversion (see Fig. 0.1).
- Mixable amounts of the chosen plasmids are gained by cultivation of the subclones and plasmid preparations. The gained plasmid stocks are equilibrated to equal concentrations before mixing.
- The plasmid stocks are mixed in unequal proportions. To gain more test mixtures from the same source the proportions are permuted. Though this is possible with many proportions we suggest to use proportions based on 2ⁿ; n ∈ [0,1,2...(cloneNumber 1)]. In the experiment we mixed the clones in the proportions 1:2:4, which resulted in eight equally distributed base compositions from 0/7 to 7/7 in steps of 1/7. Permuting the proportions allowed to generate six different mixtures¹ from the three clones which in this experiment covered many measurement points at different levels (see

¹ proportion permutations for six different mixtures of three clones: (1:2:4), (2:1:4), (1:4:2), (2:4:1), (4:1:2), (4:2:1)

- Fig. 0.2). A choice of four clones might be used for up to 24 mixtures with permutations of the proportions 1:2:4:8 leading to 16 base compositions from 0 to 1 in 1/15 steps.
- The mixtures (in the experiment six) can now be used to assess or calibrate methods which measure the base proportions at specific positions. Results from the emethod to be calibrated or assessed can be compared with expectation values based on known proportions in the test system. An example for this is given in Fig 0.2 b.

Use and advantages

Reproducibility

Once a test system like the one described is established it can easily and cheaply be reproduced with low effort and low risk of changes. More complex systems needing more preparation steps (than concentration measurement and mixing), e.g. random PCR or enzymatic preparation steps, might not be as robust as the provided system and have a high variance from batch to batch. All these characteristics make test systems based on the described method a potential commercial product: easy, reliably and cheap to produce as soon as established.

Different proportions within one mixture

Mixtures of e.g. methylated and unmethylated DNA only provide one defined base proportion to be expected after conversion, it is equal at each position. Any problems that might occur from the fact that other positions have other rates are omitted from such system. The test system described here provides different proportions at different positions within one mixture. Therefore it overcomes the problem of the other system, wherein equal proportions at all positions are used and thereby might bias measurements. In addition this method allows to generate data over a range of measurement points and not only at one defined value, therefore a single mixture can be used to assess the whole range of a measurement method.

Specialized tests based on real DNA patterns

The method allows to generate test systems providing any wanted composition of base proportion at different DNA positions whenever a needed pattern can be found in subclones derived from real samples. This allows to always choose the appropriate subclones for any analysis method the test system will be applied

to. It is e.g. possible to choose stretches that show blocks with equal base proportions at all sites of interest. This way the influence of such blocks (like local co-methylation) on measurement methods can be assessed. The fact that real sample material can be used for the initial step of subclone generation allows to easily reproduce patterns as observed in nature. E.g. for methylation analysis this offers, the opportunity to test sensitive detection methods very precisely and in detail, and allows modeling reality in a more appropriate way than by mixing DNA of 0% and 100% methylation at all positions.

Single method step assessment

The generation of mixtures of e.g. methylated and unmethylated DNA requires several steps until it can be used to assess a measurement method based on e.g. PCR products of bisulphite treated DNA. All these steps influence the real expectation values and the results. 1. the production of methylated DNA may be incomplete and introduce errors. 2. the bisulphite conversion might be incomplete 3. the amplification in the PCR might be biased or have a high variance. All these steps add to any variance and/or bias in the final measurement method to be assessed and cannot easily be separated from it. In contrast the here provided test system allows to asses measurement methods as a whole or its single steps. It therefore provides detailed information about single steps and can locate error sources more easily than methods that provide only an assessment of a whole pipeline of steps.

ABSTRACT

I here present a method that allows to assess and calibrate methods and systems that quantify base compositions at special positions in DNA. The method is characterized by using synthetic, highly reproducible test systems. Said test systems are characterized by a) being built by DNA subclone mixtures, b) providing high numbers of measuring points within one DNA subclone mixture.

The measuring points cover the range of the measurement method to be assessed or calibrated in an evenly distributed manner.

If used to assess a DNA methylation detection method the test system is able to test te outcome of single steps of said method and therefore has a huge advantage compared to methods that can only assess the outcome of multiple steps.

The method is characterized by the use of mixtures of subclones from one and the same DNA region that show base composition differences at positions of interest. The method is further characterized by taking more than two subclones, that among one another are as unequal as possible and mix them in permutations of different portions.

The test system allows to build models with patterns very close to observations in real DNA. An established system can easily be used as a standard for optimization and calibration experiments for different methods and is a potential commercial product.

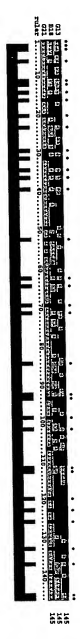


Fig. 0.1: Three final clones chosen for the mixtures, only genomic C positions and their on bisulphite treatment based equivalent (T) are shown.

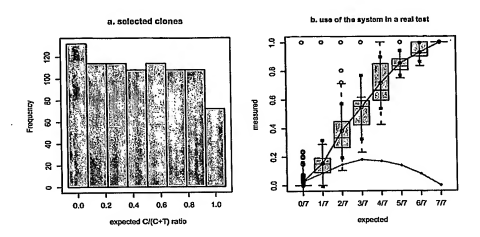


Fig. 0.2: a. Number of measuring points for different C/(C+T) ratios within all six subclone mixtures of the example. b. real calibration data based on an assessment of base ratio detection with four dye capillary sequencing.

APPENDIX: data from 96 subclones

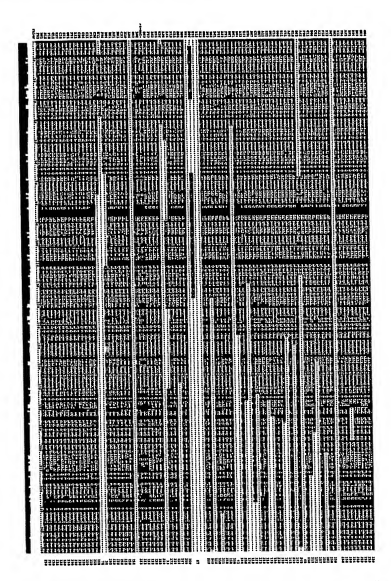


Fig. 0.3: Full sequence of clones from the initial sub-cloning step of G6e (part 1)

apigenomies

Jörn Lewin Epigenomics AG Berlin

page 8

---- AVAILABLE COPY

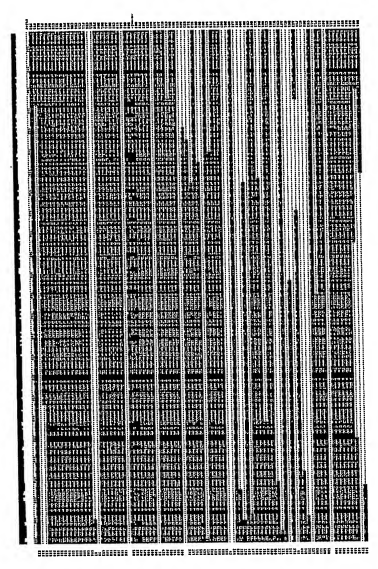


Fig. 0.4: Full sequence of clones from the initial sub-cloning step of G6e (part 2)

#pigunomies

Jörn Lewin Epigenomics AG Berlin

page 9

BEST AVAILABLE COPY

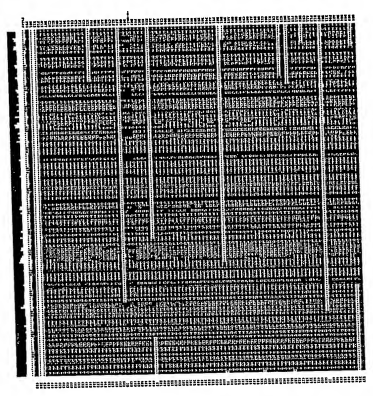


Fig. 0.5: Full sequence of clones from the initial sub-cloning step of G6e (part 3)